Light driven water oxidation by a single site cobalt salophen catalyst†

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A salophen cobalt(II) complex enables water oxidation at neutral pH in photoactivated sacrificial cycles under visible light, thus confirming the high appeal of earth abundant single site catalysis for artificial photosynthesis.

Inspired by the natural Mn₄CaO₆ oxygen evolving centre in photosystem II,1 remarkable efforts have been dedicated towards the development of multinuclear transition metal complexes enabling water oxidation for artificial photosynthesis.2,3 Multimetallic catalysts could in principle distribute the oxidation equivalents necessary for water oxidation over several metal centres, thus lowering the energy barrier of the overall catalytic process.3 However, the design of multi-metallic cores with oxygen evolving activity poses synthetic and stability hurdles.3

Noteworthily, single site metal complexes have been recently discovered, whose oxygen evolving activity offers a major opportunity to broaden the catalyst space within fundamental coordination chemistry.4 Of particular interest are the earth-abundant cobalt complexes featuring polydentate organic ligands, such as corroles, polypyridines, porphyrins, and polyamines, which have been used under dark electrocatalysis conditions,5 and in few cases also within photoactivated cycles.6 Ligand diversity is expected to play a crucial role in tuning photocatalytic water oxidation, with the urgent quest to both optimize sequential photoinduced electron transfer and facilitate the dark-phase of the catalytic mechanism under a turnover regime.

In this communication we present a cobalt(II) complex with a salophen ligand (\(N,N^-\)bis(salicylaldehyde)-1,2-phenylenediamine), CoSlp, capable of water oxidation under visible light irradiation, with Ru(bpy)₃²⁺ (bpy = 2,2'-bipyridine) as the photosensitizer and persulfate (\(S_2O_8^{2-}\)) as the sacrificial electron acceptor (Scheme 1). Combined UV-vis, dynamic light scattering (DLS) and Electron Paramagnetic Resonance (EPR) evidence identifies CoSlp as a competent oxygen evolving catalyst (OEC), enabling a two-fold photoinduced electron transfer in the ms time-frame. Our results confirm the “privileged” nature of the salophen ligand environment, readily available from simple condensation reactions, with wide applicability in different fields of catalysis, including bio-inspired oxidations.7

CoSlp is obtained by direct reaction of cobalt acetate with the salophen ligand (\(N,N^-\)-bis(salicylaldehyde)-1,2-phenylenediamine), then confirmed using ESI-MS, where a base peak at \(m/z = 374\) was observed, ascribed to the \([\text{CoSlp}H]^+\) ion (Fig. S4, ESI†). In the solid state, the cobalt ion in CoSlp displays a square planar geometry,8c while in aqueous solution it extends the coordination sphere to square pyramidal, by ligation of a water molecule in the apical position.8d Spectrophotometric titration yields a \(pK_a = 6.40\) for the aquo ligand (Fig. S3, ESI†),8 which is therefore expected to be deprotonated at neutral pH, turning into a hydroxo moiety.

Characterisation of CoSlp by cyclic voltammetry in aqueous phosphate buffer at pH 7.1 shows the onset of an intense anodic wave beginning at ca. 0.90 V (vs. Ag/AgCl) reaching a peak current of
250 μA (current density of 3.5 mA cm⁻²) at 1.35 V (Fig. S6, ESI†). The attribution of this intense wave to catalytic water oxidation was supported by the presence of a cathodic wave at −230 mV, observed only in the reverse scan, due to reduction of dioxygen formed at the working electrode (Fig. S6, ESI†). Although electrodeposited hetero-
genous cobalt oxide can possibly plague the voltammetric analysis, the electrochemical study allows for some key observations, namely: a marked difference in the CoSpl voltammogram with respect to the “ligand-free” cobalt aquo complex, known to evolve into electro-
catalyst working electrode (Fig. S6, ESI†). Although electrodeposited hetero-
genous cobalt oxide can possibly plague the voltammetric analysis, the electrochemical study allows for some key observations, namely:

– an increase of the normalized current (defined as the current divided by the square root of the scan rate) at low scan rates which confirms the presence of a rate determining chemical step preceding electron transfer (Fig. S6, ESI†),
– a very low operating overpotential, $\eta = 0.3$ V at 0.7 mA cm⁻² current density, which outperforms previously reported cobalt OECs, with overpotentials in the range 0.5–0.6 V. This latter observation is pivotal to light-driven water oxidation under homogeneous conditions, usually achieved in the presence of a photosensitizer and a sacrificial electron acceptor, respectively, Ru(bpy)³⁺ and $S_2O_8^{2-}$ in the present case (eqn (1)–(5) in Scheme 1). In such a system, the photogenerated oxidant is the Ru(bpy)³⁺ species, produced by reaction of the excited state of Ru(bpy)³⁺ with the persulfate anion (eqn (1)–(3)). The Ru(bpy)³⁺ then oxidizes the catalyst CoSp to an activated form, generally indicated as CoSp³⁺ (eqn (4)), which is capable of oxidizing water to oxygen restoring the initial state of the catalyst (eqn (5)).

The photosynthetic activity of CoSp (15–125 μM) was confirmed by oxygen evolution upon illumination of a 20 mM phosphate buffered aqueous solution (pH 7) containing 1 mM Ru(bpy)³⁺ and 5 mM $S_2O_8^{2-}$ (Fig. 1a). An initial lag-time is observed in the kinetic traces, due to accumulation of a steady-state concentration of the photogenerated intermediates, as it is eliminated by increasing the irradiation flux. The maximum turnover rate and the total amount of oxygen produced depend on the catalyst concentration, showing a saturation-inhibition behavior for [CoSp] > 100 μM (Fig. 1a and Table S1, ESI†). The oxygen production levels off after ca. 2 hours due to Ru(bpy)³⁺ degradation, as revealed by the partial bleaching of the absorption at 450 nm (Fig. S7, ESI†). In this timeframe, CoSp operates for up to 17 turnovers. The most significant parameter to describe the performance of a photoactivated system is the quantum yield ($\Phi_{O_2}$), defined as the ratio between the oxygen produced and the photons absorbed by the system (eqn (6)).

\[ \Phi_{O_2} = \frac{O_2 \text{ produced}}{\text{absorbed photons}} \]  

Under the explored conditions, $\Phi_{O_2}$ was found in the range 0.048–0.079, corresponding to 9.6–15.8% efficiency in photon to $O_2$ conversion, with a bell-shaped profile depending on CoSp concentration (Fig. 1b). A similar bell-shaped profile was already observed with a tetracobalt cubane catalyst, and abatement of the rate of $O_2$ production at >100 μM cobalt concentration was indeed noted also with other molecular precatalysts. This behavior is related to competitive unproductive routes amplified at high catalyst concentration, and likely identifiable with quenching of the Ru(bpy)³⁺ excited state by CoSp and by its intermediates involved in the catalytic cycle.

In photoactivated cycles devoted to oxygen production, a critical step that often determines the overall efficiency is the electron transfer rate from the catalyst to the oxidized photosensitizer (eqn (4) in Scheme 1). This can be conveniently investigated by performing laser flash photolysis experiments, where a suitable concentration of Ru(bpy)³⁺ is generated in a <10 ns timeframe by 355 nm laser activated reaction of Ru(bpy)³⁺ with $S_2O_8^{2-}$ (eqn (1)–(3)), while the following reaction of Ru(bpy)³⁺ with CoSp is monitored in a ms timescale by the recovery of the absorption at 450 nm, due to Ru(bpy)³⁺ regeneration (Fig. 2). As shown in Fig. 2a, the rate of recovery of the absorption of Ru(bpy)³⁺ at 450 nm depends on CoSp concentration, and assuming pseudo-first order kinetic conditions, a bimolecular rate constant of $1.12 \times 10^8$ M⁻¹ s⁻¹ is obtained for the first electron transfer event (Fig. S8, ESI†), likely involving the formation of a Co(m) intermediate. This value is one order of magnitude higher with respect to a tetracobalt cubane catalyst, explored under the same conditions.

Moreover, examination of the process with sub-stoichiometric catalyst solutions shows that each CoSp is able to scavenge ca. two Ru(bpy)³⁺ within a time window of 40 ms (Fig. 2b). This suggests the fast formation of a formal Co(m) derivative by two consecutive one-electron oxidation of CoSp by Ru(bpy)³⁺ under irradiation conditions. A possible mechanistic scenario foresees the involvement of a photogenerated Co(m)-oxo intermediate, undergoing a rate determining nucleophilic attack by a water molecule. A new oxygen–oxygen bond is then formed, within the structural motif of Co(m)-hydroperoxide (Scheme 2); such an intermediate could finally release $O_2$ upon further oxidation.

As a final remark, stability is a major requirement in order to consider a catalyst for potential application in regenerative oxygen evolving photoanodes. In the present case, UV-vis experiments reveal unchanged absorption spectra over several hours of CoSp in 20 mM aqueous phosphate buffer (pH 7), also in the presence of the persulfate anion. Under photocatalytic conditions, dynamic light scattering analysis of the reaction mixtures yields no evidence of any
dispersion, suggesting the absence of Co-oxide colloids. Moreover, the reaction mixture has been analysed using X-band continuous-wave (CW) EPR spectroscopy (Fig. 3). When frozen in a CH$_3$Cl$_2$ : CH$_3$OH 9 : 1 matrix, CoSlp shows an orthorhombic EPR spectrum at 10 K with $g_1 = 7.90$, $g_2 = 1.90$ and $g_3 = 1.85$, where the signal due to the highest $g$ component shows an eight-line cobalt-hyperfine splitting ($A_1 = 1100$ MHz, Fig. 3a). According to spectral simulation this is consistent with an $S = 3/2$ species, as expected for a $d^7$ high spin configuration of the Co(n) ion.\textsuperscript{16,16} In the aqueous phase (CH$_3$CN : phosphate buffer 1 : 1), the EPR spectrum is characterised by the presence of a featureless and nearly axial signal with $g$ values of $g_1 = 4.95$, $g_2 = 3.75$ and $g_3 = 2.05$ (black trace in Fig. 3b), which is still assigned to an $S = 3/2$ Co(n) species.\textsuperscript{17} The occurrence of a high-spin Co(n) signal is likely due to strong axial interactions and/or distortions of the planar structure of the complex in both media.\textsuperscript{16,16,16}\textsuperscript{18} Under photocatalytic conditions, the EPR spectrum obtained for CoSlp does not undergo major changes even after prolonged illumination in the presence of Ru(bpy)$_3^{2+}$ and S$_2$O$_8^{2-}$ (blue trace in Fig. 3b). No signals of free Co(n) ions or Co(n) oxides are detectable,\textsuperscript{16,16,19} thus confirming the resting form of CoSlp and its stability under the operating conditions.

In conclusion, we have presented a novel molecular water oxidation catalyst, based on a single site cobalt salophen complex. This species catalyses water oxidation in neutral aqueous medium electrochemically and in a photocatalysed system, showing very high electron transfer rates to the photo-generated oxidant Ru(bpy)$_3^{2+}$. Structure reactivity studies,\textsuperscript{20} by introducing substituents in the aromatic rings of the catalyst, will be addressed to optimize catalyst performance.

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Notes and references

† The residual presence of an anodic wave, with ca. 30% intensity of the pristine curve, in a CoSlp-free solution, with an unpolished working electrode from a previous scan indicates the possible absorption-deposition of active material onto the working electrode.\textsuperscript{12,12} § In Ru(bpy)$_3^{2+}$–S$_2$O$_8^{2-}$ systems, the maximum value of $\phi_O$ is indeed 0.5, since two Ru(bpy)$_3^{2+}$ are possibly generated upon absorption of one photon (eqn (1)–(3)), and at least four Ru(bpy)$_3^{2+}$ are required to produce one O$_2$ molecule.\textsuperscript{1} Reported quantum efficiencies for a single site Co precatalyst reach up to 30%.\textsuperscript{12,13} ¶ The decrease in the initial AOD observed upon CoSlp addition is mainly attributed to laser absorption by CoSlp and quenching of “Ru(bpy)$_3^{2+}$” by the cobalt catalyst (Fig. S9, ESI†).

| Co(n)-oxo species may display a spin distribution on the oxygen atom, and therefore should be better described as Co(μ-O)–O$^-$ (Co-oxyl).\textsuperscript{15} |


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